

H58271, H16122, W77956, AA193332, AA323923, AA370209,
 AA296758, W72757, AA093971, AA385544, AA386175, AA165402,
 AW085713, H42806, AA093977, AI161152, AA370011, AI671702,
 R71215, AA885343, T79297, AI814869, R81567, AI082713, N29615,
 AW087726, AW075710, AI952608, AI818073, AI784445, AI432812,
 AI375568, AI372904, AI364106, AI143379, AA993074, AA953985,
 AA862385, AA761084, AA576229, AA569223, AA463198, AA452117,
 AA416877, AA074872, W16851, W04568, N40176, AW068354,
 AA857004, H58663, H15819, AW264944, AI923965, AI692214,
 AI475321, AI435987, AA961068, AA206059, AI469161, T84789,
 AA507257, AA707515, AA132458, AA179262, T79211, W31505,
 N25699, T99574, T99363, AI751598, AA713668, T91119, AW105515,
 AA370208, AI422128, R81568, AI038899, AI971847, AI540650,
 AI826106, AA885960, R56263, AA825431, T99147, D31503 and
 AF049564, or

d) a complementary sequence to the sequences of a) and/or b).

A¹
 2. Nucleic acid according to claim 1, which includes a protein-coding segment comprising [of preferably] at least 30 nucleotides of the nucleotide sequence shown in Fig. 1.

A²
 4. Modified nucleic acid or nucleic acid analogue, which includes a nucleotide sequence according to [one of the claims 1 to 3] claim 1.

5. Recombinant vector, which includes at least one copy of a nucleic acid according to [one of the claims 1 to 3] claim 1 or a section thereof.

A³
 7. A transformed cell, non-human transgenic organism, or animal model comprising [With] a nucleic acid according to [one of the claims 1 to 3] claim 1 or a vector according to claim 5 [or 6 transformed cell, a corresponding non-human transgenic organism or animal models], which stably produce (knock-in) the product of the nucleic acid according to [one of the claims 1 to 3] claim 1 or whose corresponding natural gene was destroyed deliberately (knock-out).

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8. Polypeptide or a salt thereof, which is coded by a nucleic acid according to [one of the claims 1 to 3] claim 1.
10. Fragment of the polypeptide according to claim[s] 8 [or 9] with at least 100 amino acids or salts thereof.
11. Modified polypeptide, which includes an amino acid sequence according to claim[s] 8 [or 9].
12. Method[s] for the synthesis of the polypeptide according to claim 8 [or 9], which includes the cultivation of cells according to claim 7 [as well as] and the isolation of the polypeptide according to claim 8 [or 9].
13. A method for producing an antibody against the polypeptide of claim 8, comprising contacting an antibody-producing cell with [Use of] a polypeptide according to claim 8 [or 9] or [of] fragments of this polypeptide as an immunogen [for the production of antibodies].
14. Antibodies against a polypeptide according to claim 8 [or 9].
15. Method for the identification of effectors of a protein according to claim 8 [or 9], with the help of which various potential effector substances can be tested on cells, which express the protein.
16. Pharmaceutical composition, which includes as an active component
 - a nucleic acid according to [one of the claims 1 to 4] claim 1,
 - b) a vector according to claim 5 [or 6],
 - c) a cell according to claim 7,
 - d) a polypeptide according to claim 8, [9,]10 or 11,
 - e) an antibody according to claim 14

and which contains the pharmaceutically usual carrier, auxiliary and/or additive substances.

17. A method of diagnosing a disease [Use of a composition according to claim 16 for diagnosis of diseases,] which [are] is associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression, or a predisposition to such a disease[s] comprising the use of a composition according to claim 16.

18. A method of diagnosing a disease [Use of a pharmaceutical composition for diagnosis of diseases] which [are] is associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression, or a predisposition to such a disease[s], comprising the use of a composition which contains as an active component
a) an EST sequence according to claim 1c,
b) a recombinant vector which includes at least one copy of the EST sequences mentioned above,
c) a recombinant vector according to b) which enables the expression of the nucleic acid in a suitable host cell,
d) a cell according to claim 7, whereas the nucleic acid consists of one of the EST sequences mentioned above,
e) a polypeptide being coded by one of the EST sequences mentioned above or a salt thereof or,
f) a polypeptide according to e) which exhibits the amino acid sequence shown in Fig.2 or a homology of more than 60% with the amino acid sequence shown in Fig.2 or a salt thereof,
g) a fragment of the polypeptide according to e) or f) with at least 100 amino acids or a salt thereof,
h) a modified polypeptide which includes an amino acid sequence according to e) or f),
i) an antibody against a polypeptide according to e) or f)
and which contains pharmaceutically usual carrier, auxiliary and/or additive substances.

19. A method for treating or preventing a disease [Use of a composition according to claim 16 for the therapy or prevention of diseases,] which [are] is associated with DNA repair

defects, cell cycle disorders, cytopenia, tumor genesis and/or tumor progression,
comprising administering a composition of claim 16.

20. A method for treating or preventing a disease [Use of a pharmaceutical composition according to claim 18 for the therapy or prevention of diseases.] which [are] is associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression, comprising administering a composition of claim 18.

21. The method of claim 19, wherein said treating or preventing is carried out by [Use of a composition according to claim 16 for a] gene therapy [of diseases, which are associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression].

22. The method of claim 20, wherein said treating or preventing is carried out by [Use of a pharmaceutical composition according to claim 18] for gene therapy [of diseases which are associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression].

23. Method[s] for diagnosing diseases, which are associated with DNA repair defects, cell c cycle disorders, cytopenia, tumorigenesis and/or tumor progression or a predisposition to such diseases, during which a patient or a sample from the patient is brought in contact with a composition according to claim 16 and the nucleotide sequence and/or the expression of a nucleic acid according to claim 1 is determined.

24. Method[s] for the therapy or prevention of diseases, which are associated with DNA repair defects, cell cycle disorders, cytopenia, tumorigenesis and/or tumor progression, during which a patient is administered a composition according to claim 16, which contains the active components in an amount effective against the disease.

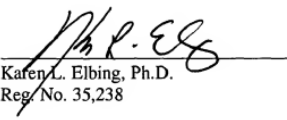
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Respectfully submitted,

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Karen L. Elbing, Ph.D.
Reg. No. 35,238

Clark & Elbing LLP
176 Federal Street
Boston, MA 02110-2214
Telephone: 617-428-0200
Facsimile: 617-428-7045

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